

Fig. 1

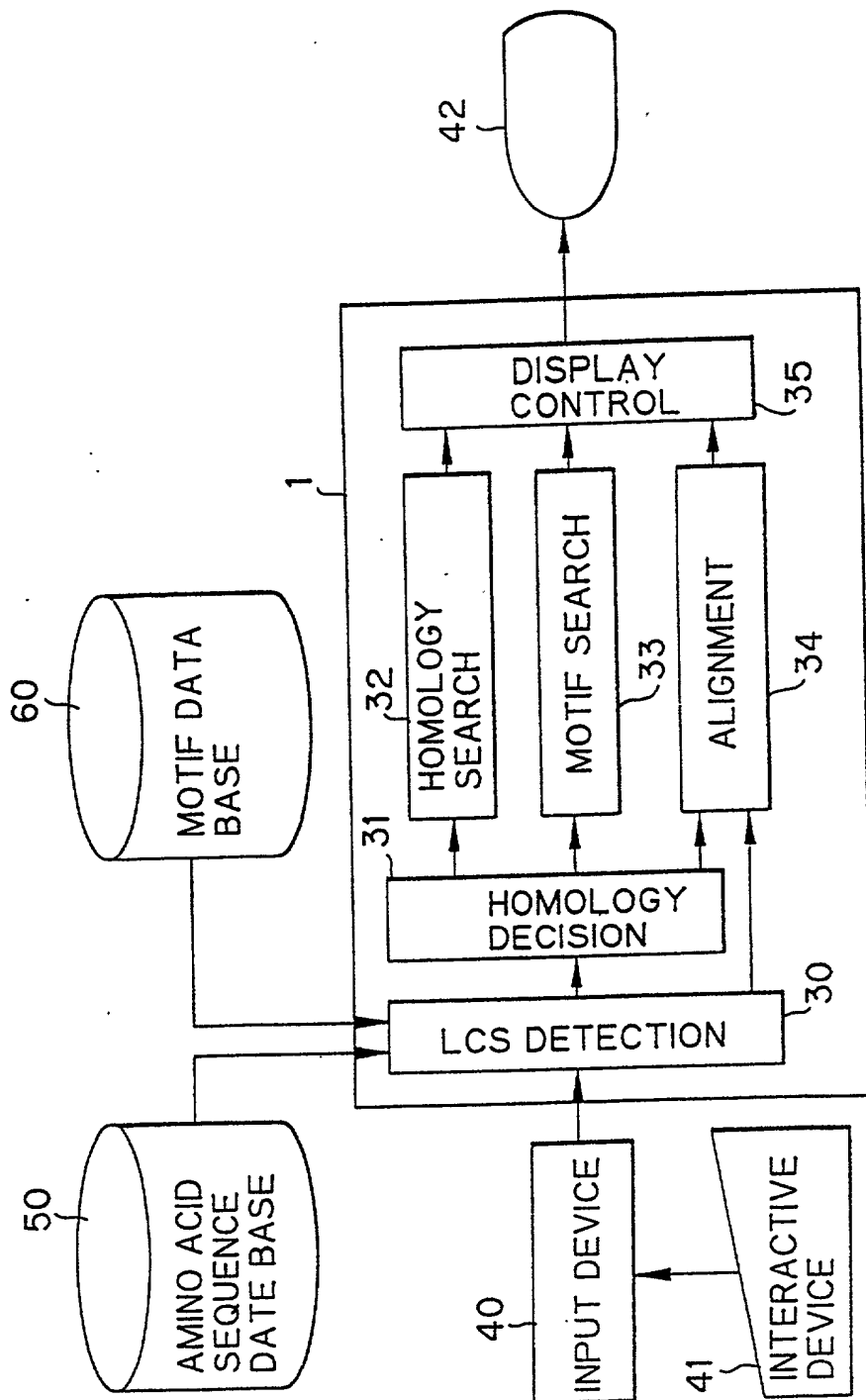


Fig. 2

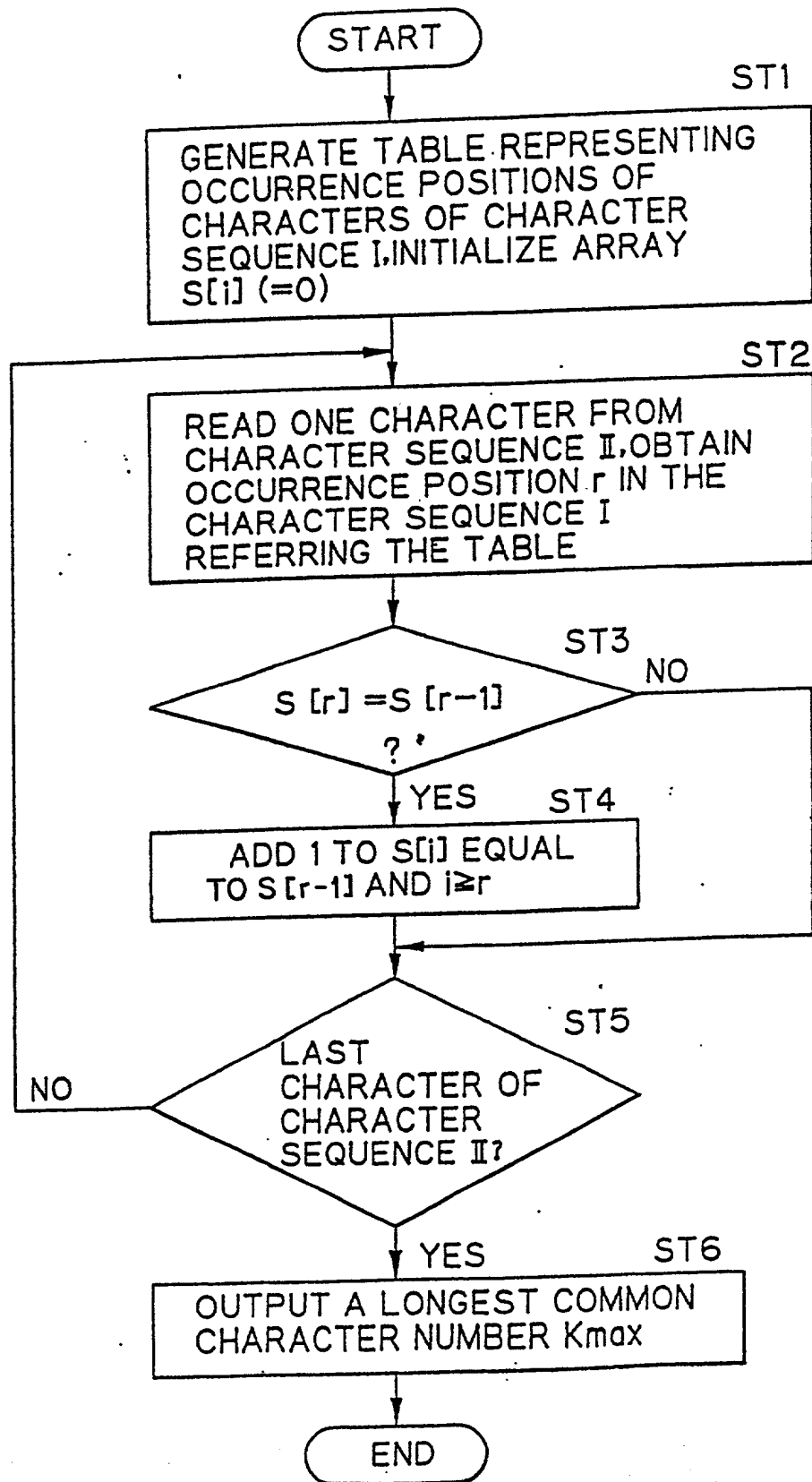


Fig. 3

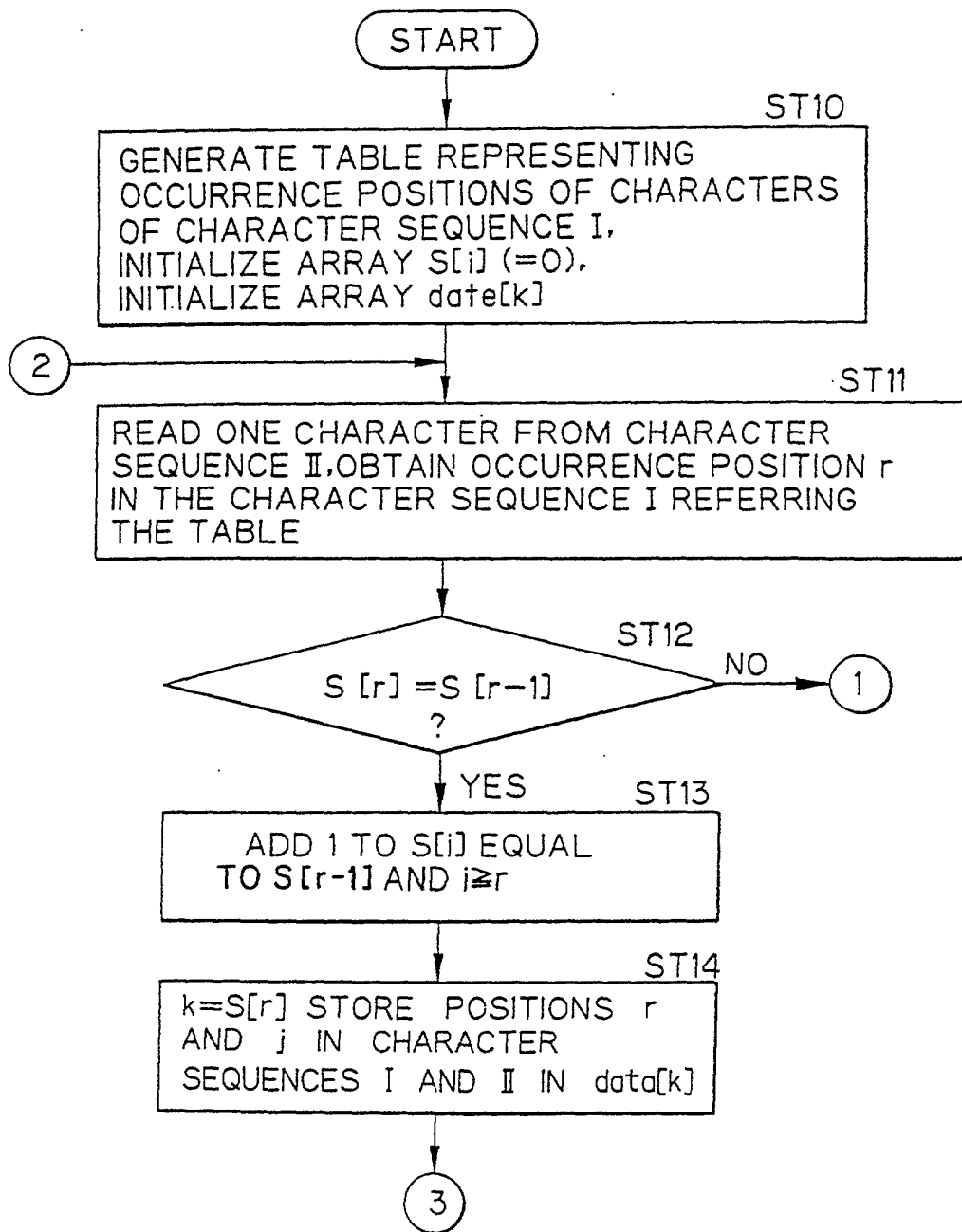
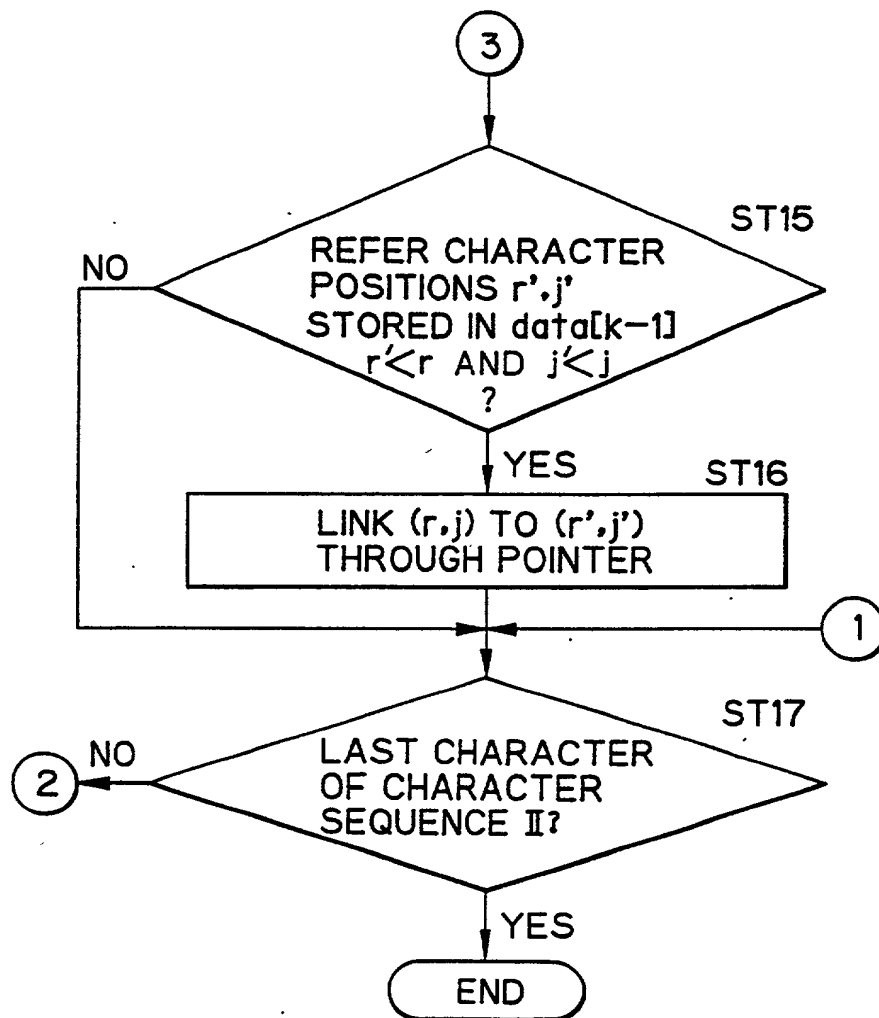


Fig. 4



*Fig. 5*

CHARACTER SEQUENCE I="ABCB DAB"

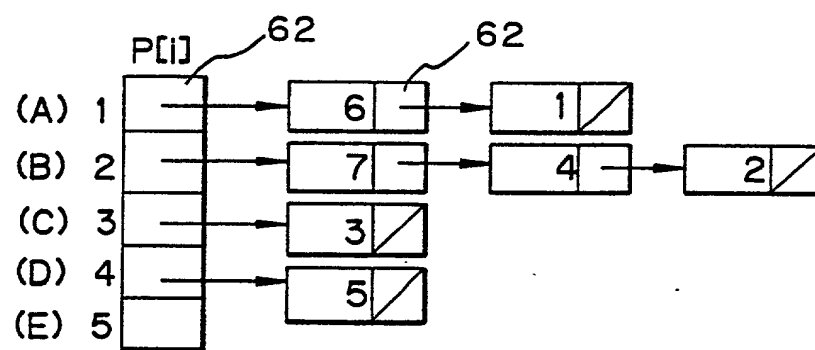
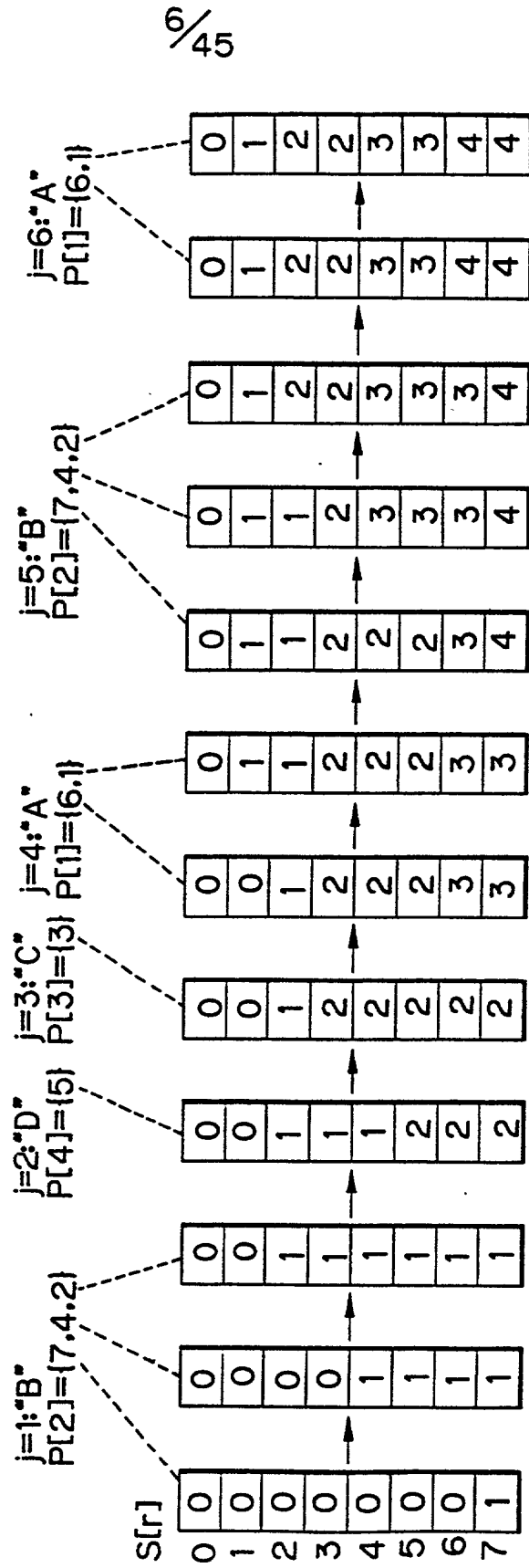


Fig. 6

CHARACTER SEQUENCE II = "BDCABA"





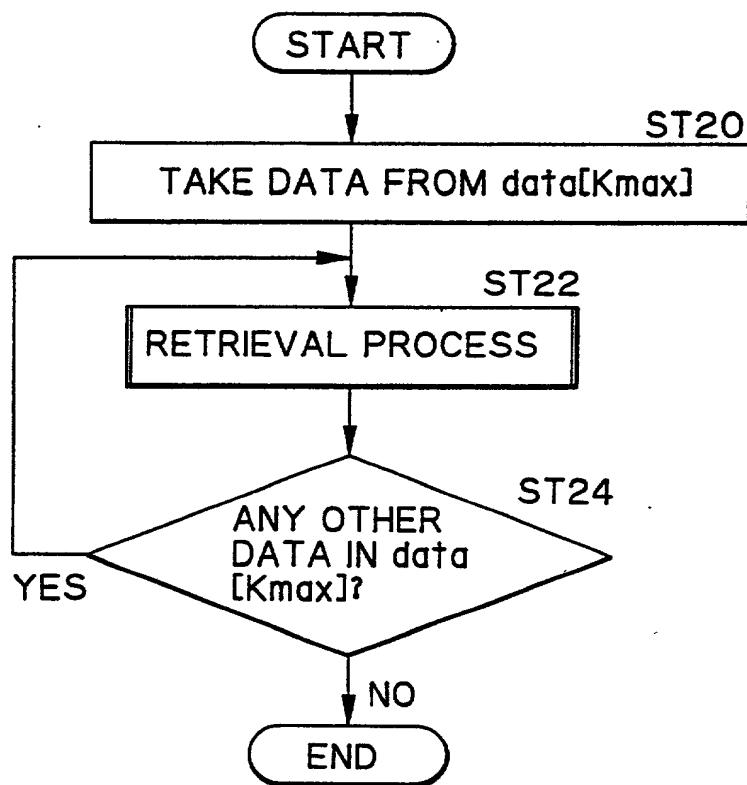
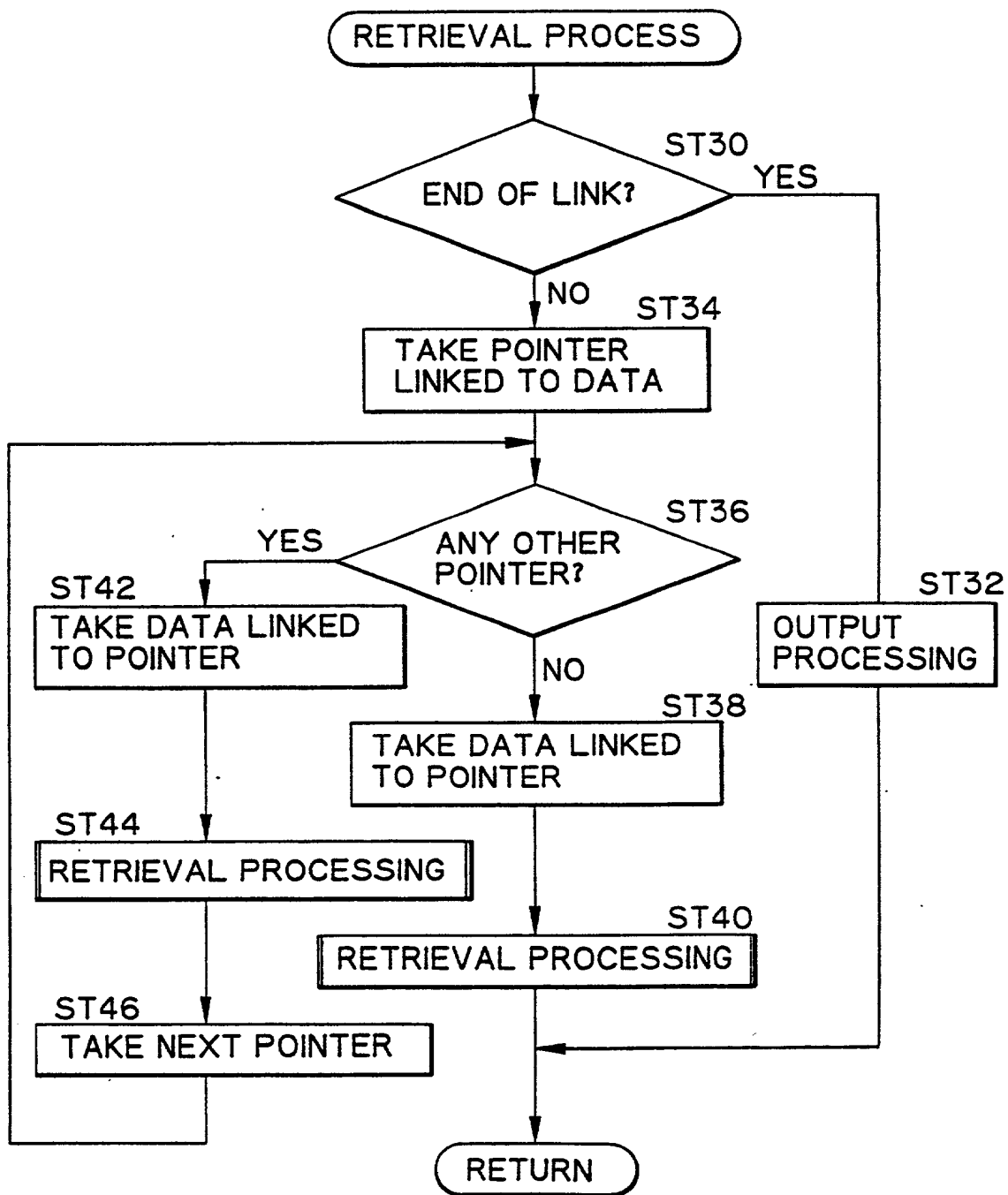
*Fig. 8*



Fig. 9



**Fig. 10**

human : GDVEKGKKIFIMKCSQCHTVEGGKHKTGPNLHGLFGRK  
bacterium : EGDAAGEKVSCKCLACHTFDQGGANKVGNPNLFGVF

LCS : GD{x3.3}G{x0.1}K{x0.2}K{x4.0}KC{x2.2}CHT{x3.3}GG{x2.2}K  
GD{x1.4}E{x0.2}K{x0.2}K{x0.4}KC{x2.2}CHT{x3.3}GG{x2.2}K

homology : 47%

**Fig. 11**

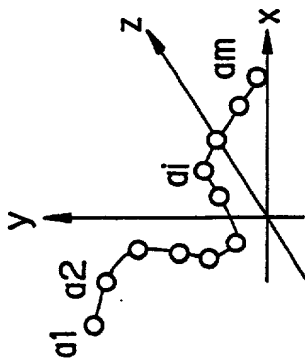
Rat : MSLAILRVIRLVRVFRIFKLSRHSKGLQILGRTLKASMRELGLLFFIGVV

**leucinzip.** L{6}L{6}L{6}L{6}L

*Fig. 12*

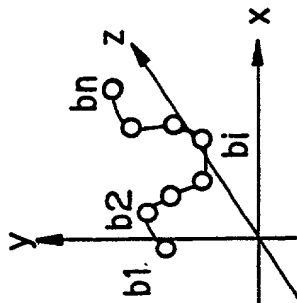
human : GDVEK G K KIFIMKCSQCHTVEKGG KHK TGP NLHGLFGRK ...  
bacterium : E GDAAAGEKVSK KCLACHTFDQGGANKV GP NPN LFGVF...

Fig. 13 A



$$A = \{a_1, a_2, \dots, a_i, \dots, a_m\}$$

Fig. 13 B



$$B = \{b_1, b_2, \dots, b_j, \dots, b_n\}$$

Fig. 13 C

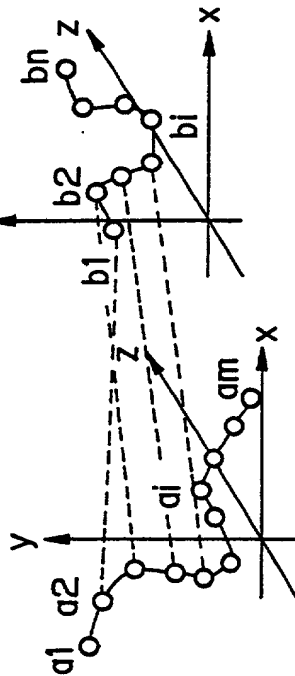


Fig. 13 D

$$r.m.s.d = \frac{\sqrt{\sum_{k=1}^n w_k (U_{bk} - a_k)^2}}{n}$$



Fig. 15

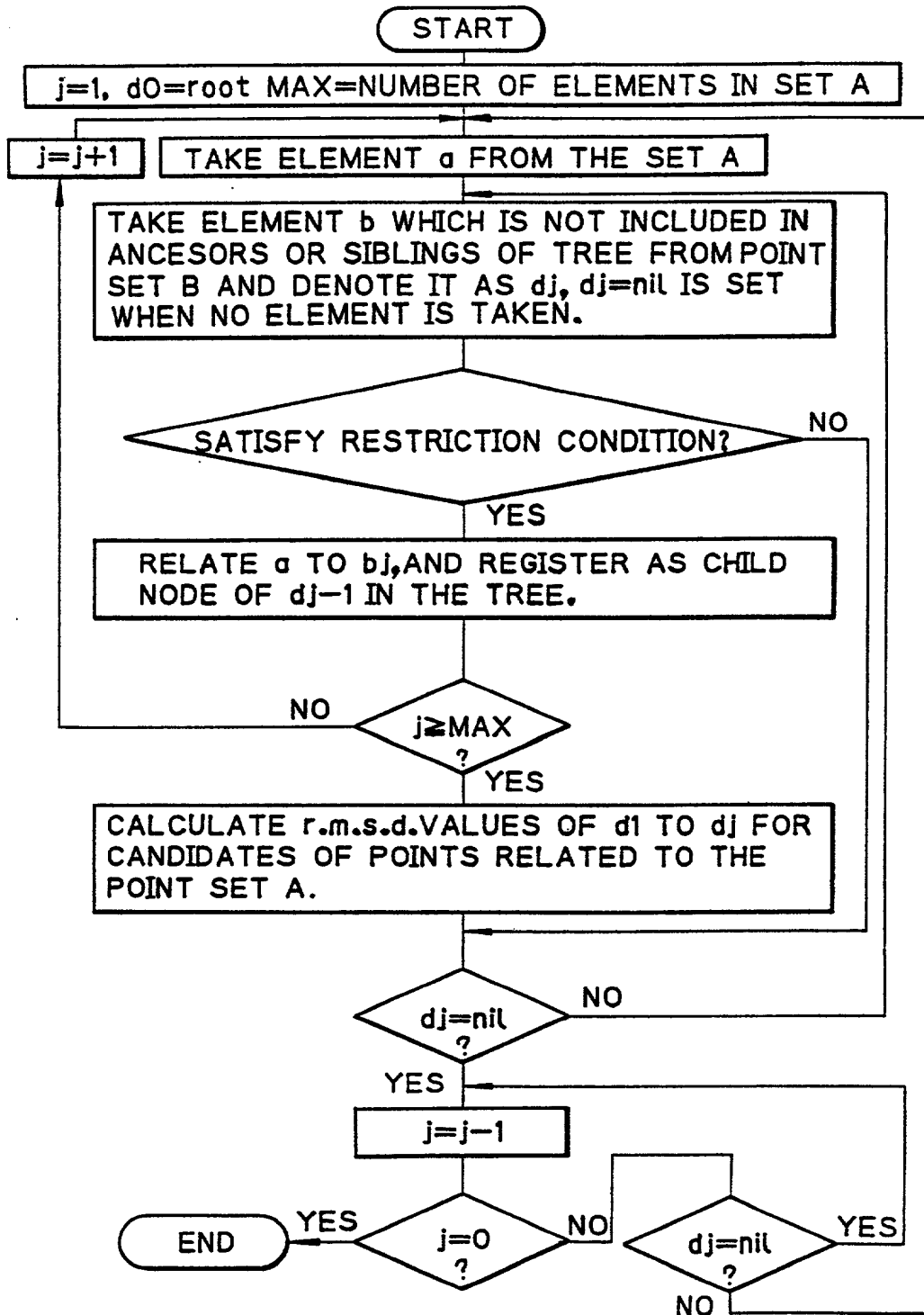


Fig. 14 A

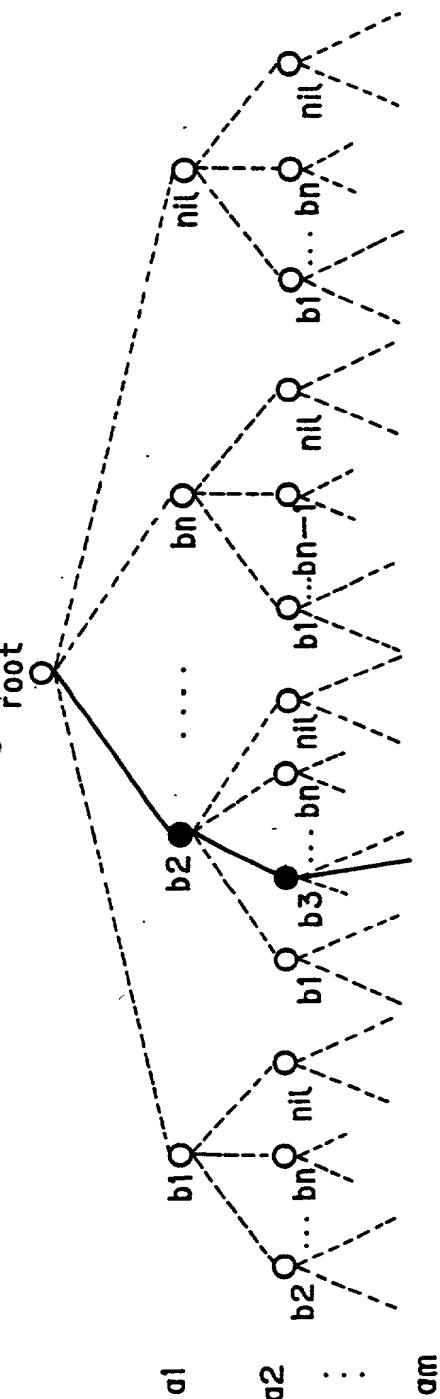


Fig. 14 B

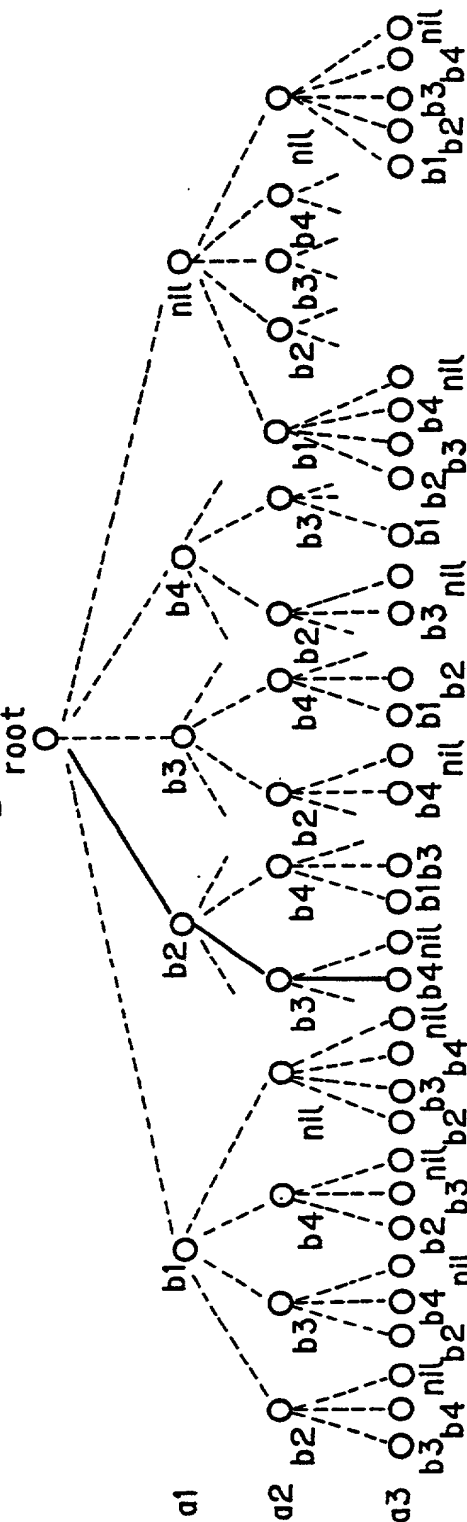


Fig. 16 A

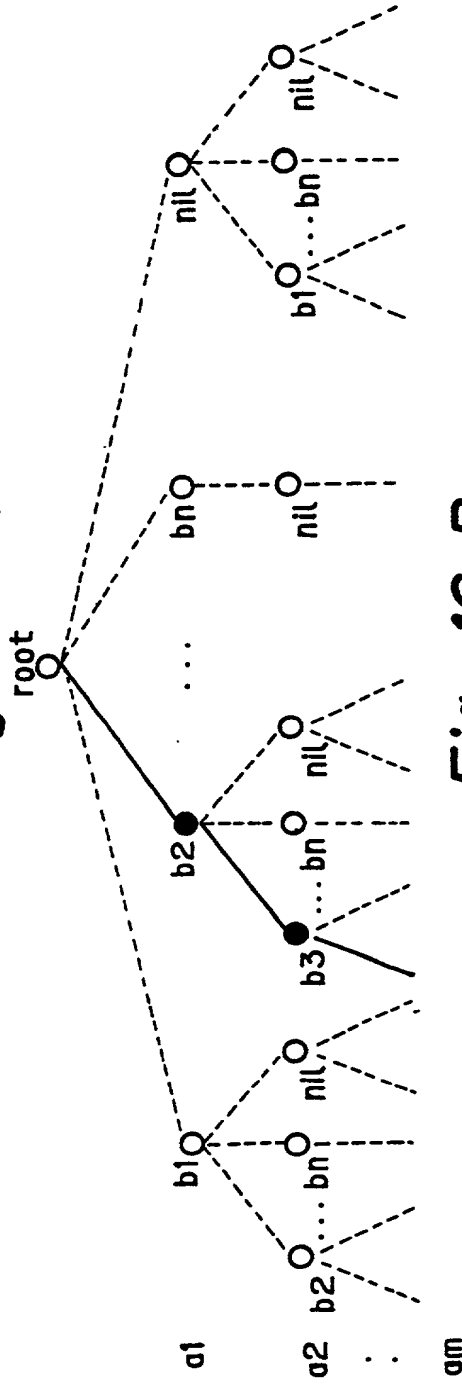


Fig. 16 B

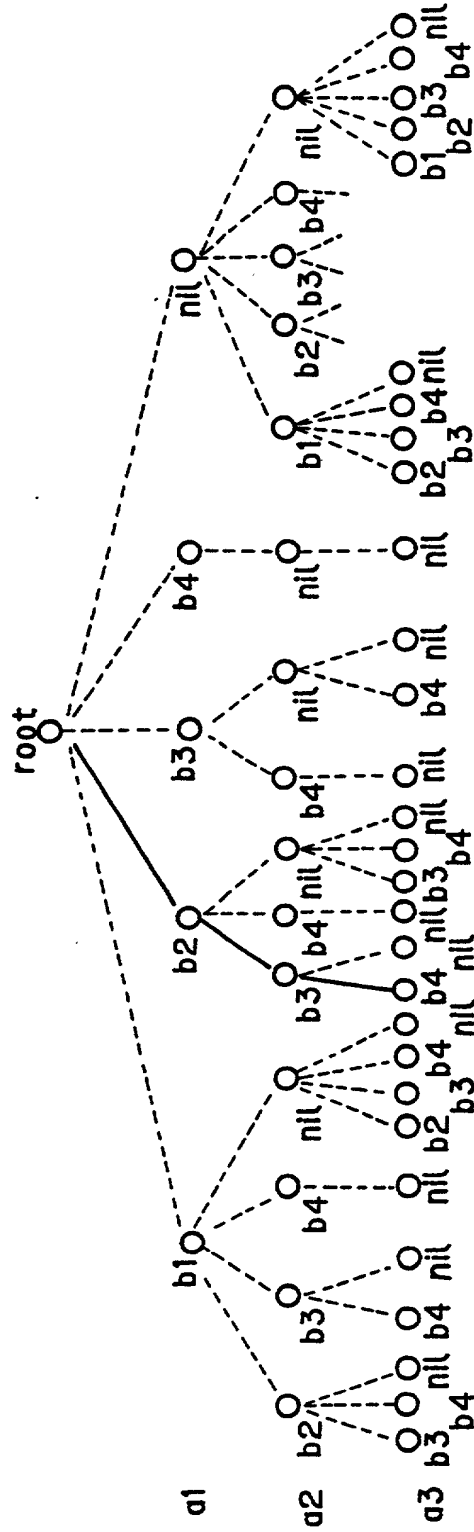
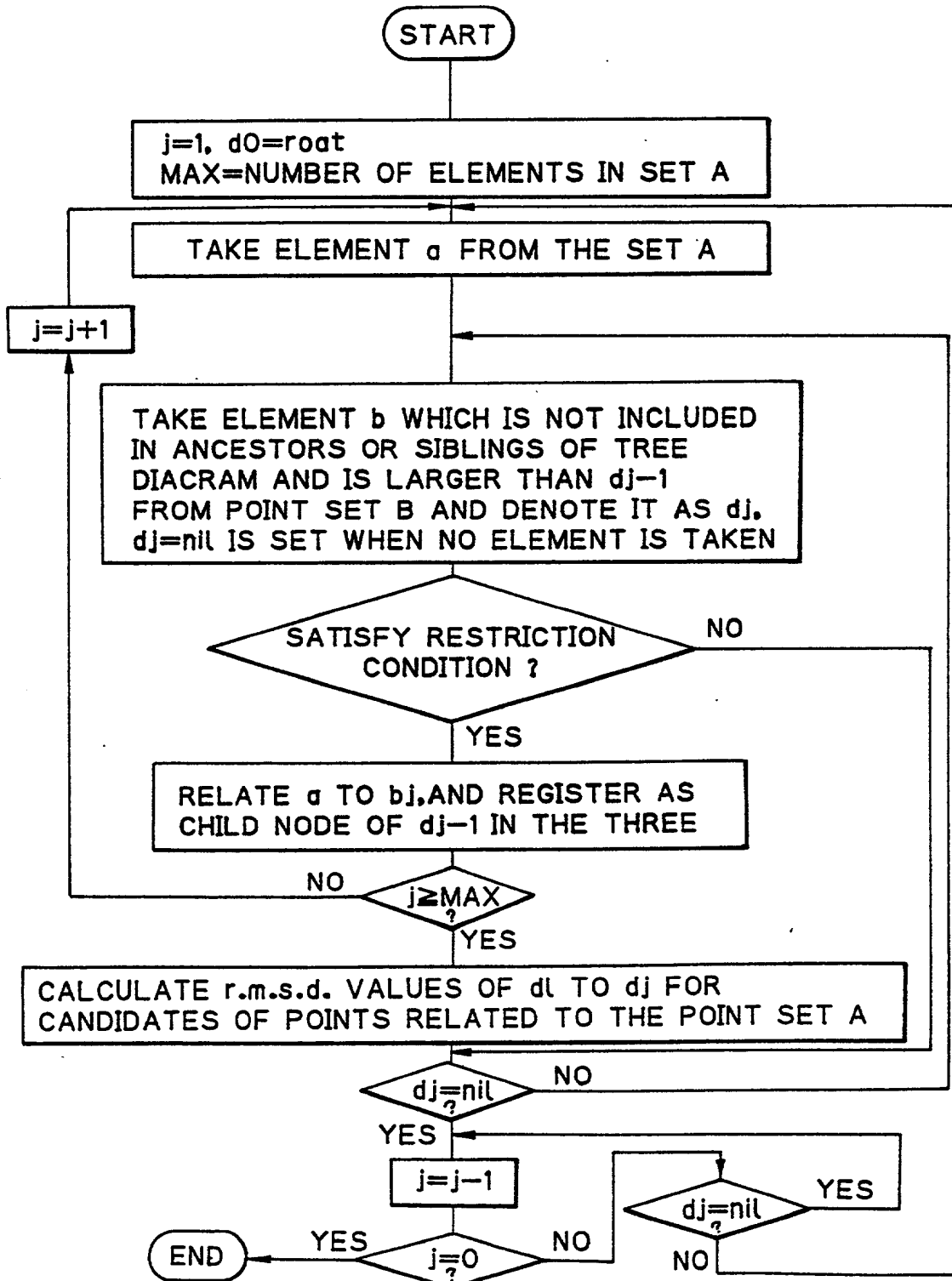
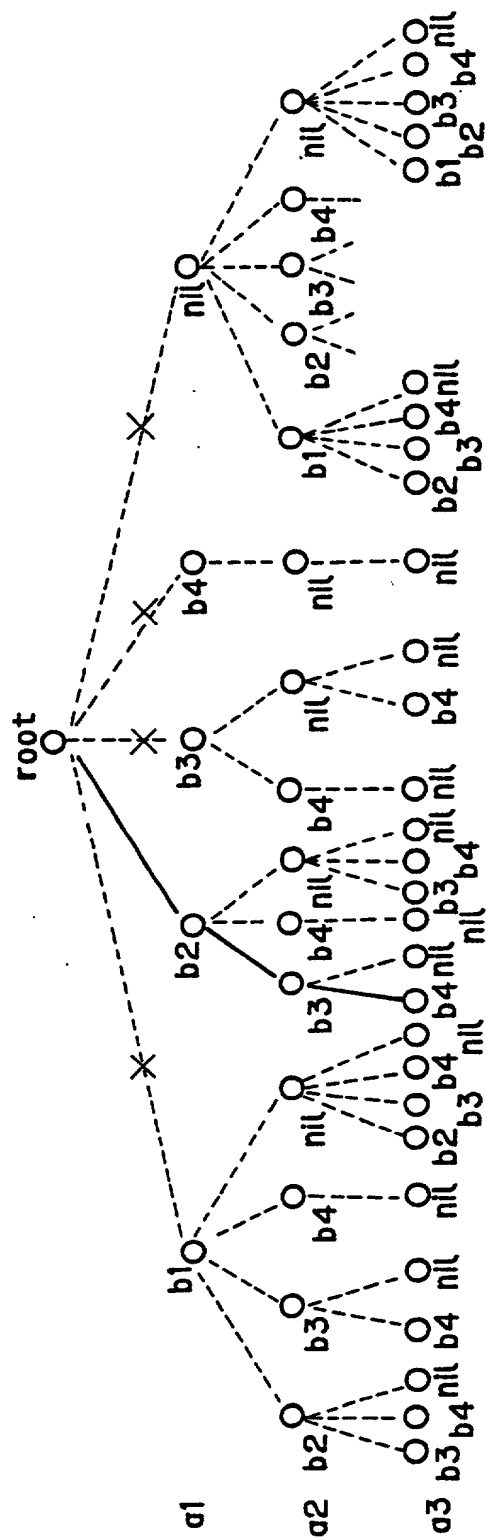




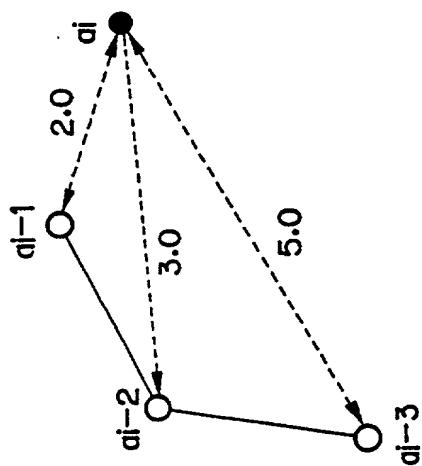
Fig. 17



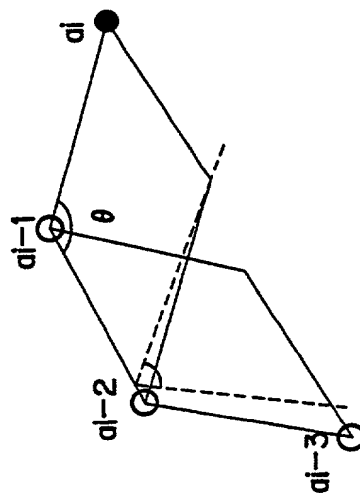
**Fig. 18**



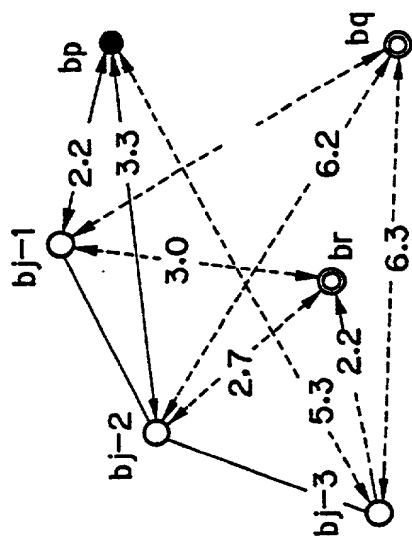
**Fig. 19 A**



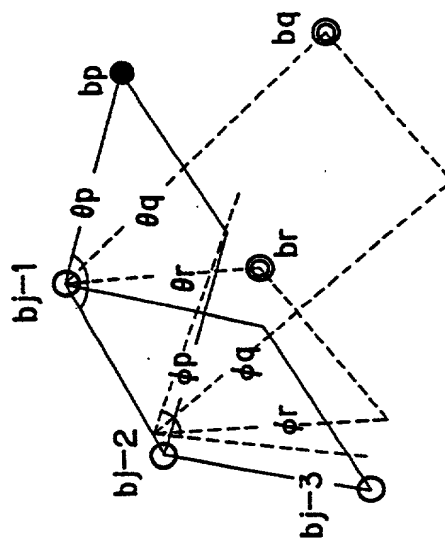
**Fig. 20 A**



**Fig. 19 B**

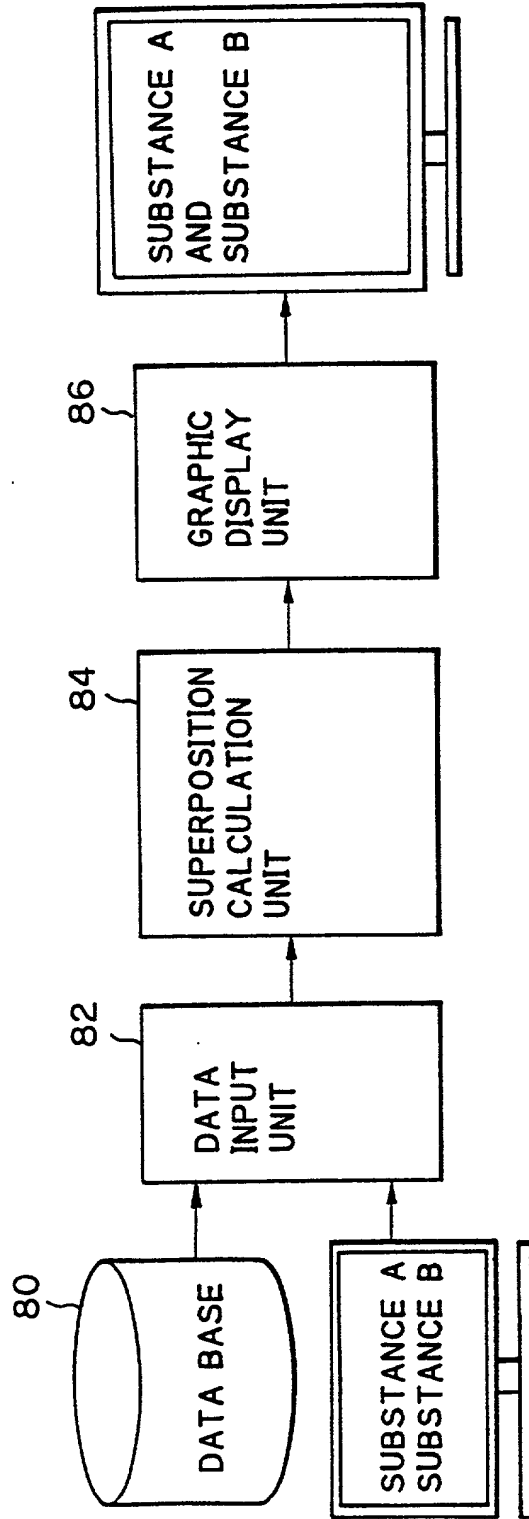


**Fig. 20 B**



The diagram illustrates a search tree for a 4-bit vector. The root node branches into three main paths: a1, a2, and a3. Each path further branches into b1, b2, b3, and b4, which then branch into nil or further b nodes. The diagram shows a search process where many branches are pruned, indicated by 'X' marks on the edges. The final nodes are labeled with bit patterns like b1 b2 b3 b4, nil, etc.

Fig. 22



**Fig. 23 A**

1	TEEQIAEFKE	AFSLFDKDG
21	GTITTKELGT	VMRSLGQNPT
41	EAELQDMINE	VDADGNGTID
61	FPEFLTMMAR	KMKDTDSEEE
81	IREAFRVFDK	DGNGYISAAE
101	LRHVMTNLGE	KLTDDEEVDEM
121	IREANIDGDG	QVNYEEFVQM
141	MTA	

AMINO ACID SEQUENCE OF CALMODULIN  
(EXCERPT FROM PDB)

**Fig. 23 B**

1	AMDQQAEARA	FLSEEMIAEF
21	KAAFDMFAD	GGGDISTKEL
41	GTVMRMLGQN	PTKEELDAAI
61	EEVDEEDGSGT	IDFEEFLVM
81	VRQMKEDAKG	KSEEELADCF
101	RIFDKNADGF	IDIEELGEIL
121	RATGEHVTEE	DIEDLMKDSD
141	KNNDGRIDFD	EFLKMMEGVQ
161		

AMINO ACID SEQUENCE OF TROPONIN C  
(EXCERPT FROM PDB)

Fig. 24 A

CALMODULIN

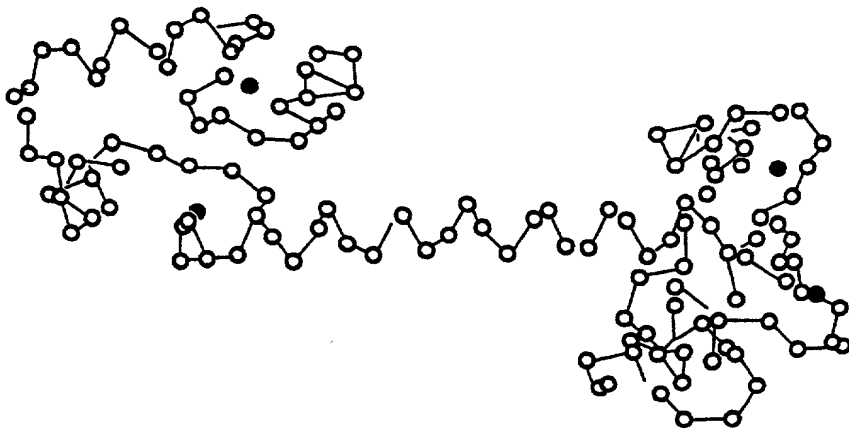
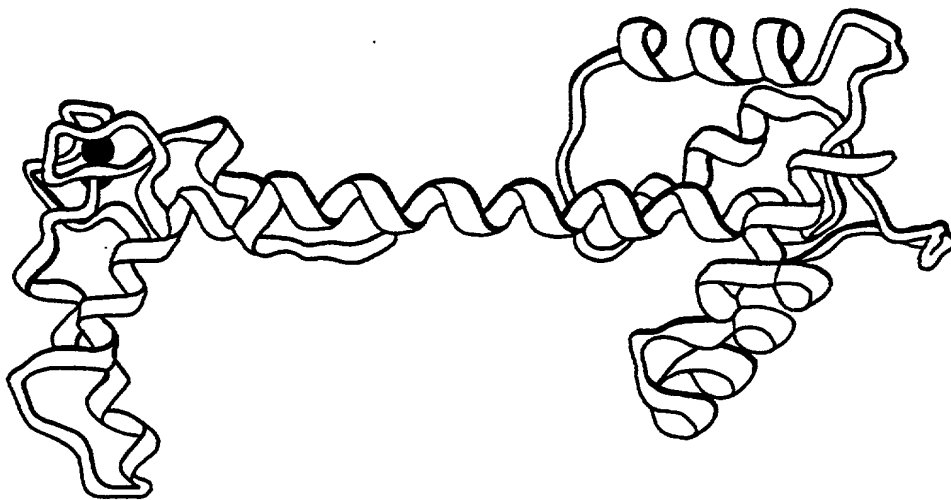


Fig. 24 B

TROPONIN C



**Fig. 25**

Probe site = 81-108 in Calmodulin																			
96	97	98	99	100	101	102	103	104	105	106	107	108	109	110					
L	A	D	C	F	R	I	F	D	K	N	A	D	G	F	< target	>			
I	R	E	A	F	R	V	F	D	K	D	G	N	G	Y	< probe	>			
111	112	113	114	115	116	117	118	119	120	121	122	123							
I	D	I	E	E	E	L	G	E	I	L	R	A	T	< target	>				
I	S	A	A	E	E	L	R	H	V	M	T	N	L	< probe	>				

**rmsd = 0.567034**



**Fig. 26**

Probe site = 81-108 and 117-143 in Calmodulin

96	97	98	99	100	101	102	103	104	105	106	107	108	109	110	
L	A	D	C	F	R	I	F	D	K	N	A	D	G	F	< target >
I	R	E	A	F	R	V	F	D	K	D	G	N	G	Y	< probe >
111	112	113	114	115	116	117	118	119	120	121	122	123			
I	D	I	E	E	L	G	E	I	L	R	A	T			< target >
I	S	A	A	E	L	R	H	V	M	T	N	L			< probe >
132	133	134	135	136	137	138	139	140	141	142	143	144	145		
I	E	D	L	M	K	D	S	D	K	N	N	D	G		< target >
V	D	E	M	I	R	E	A	N	I	D	G	D	G		< probe >
146	147	148	149	150	151	152	153	154	155	156	157	158			
R	I	D	F	D	E	F	L	K	M	M	E	G			< target >
Q	V	N	Y	E	E	F	V	Q	M	M	T	A			< probe >

rmsd = 0.823665

Fig. 27

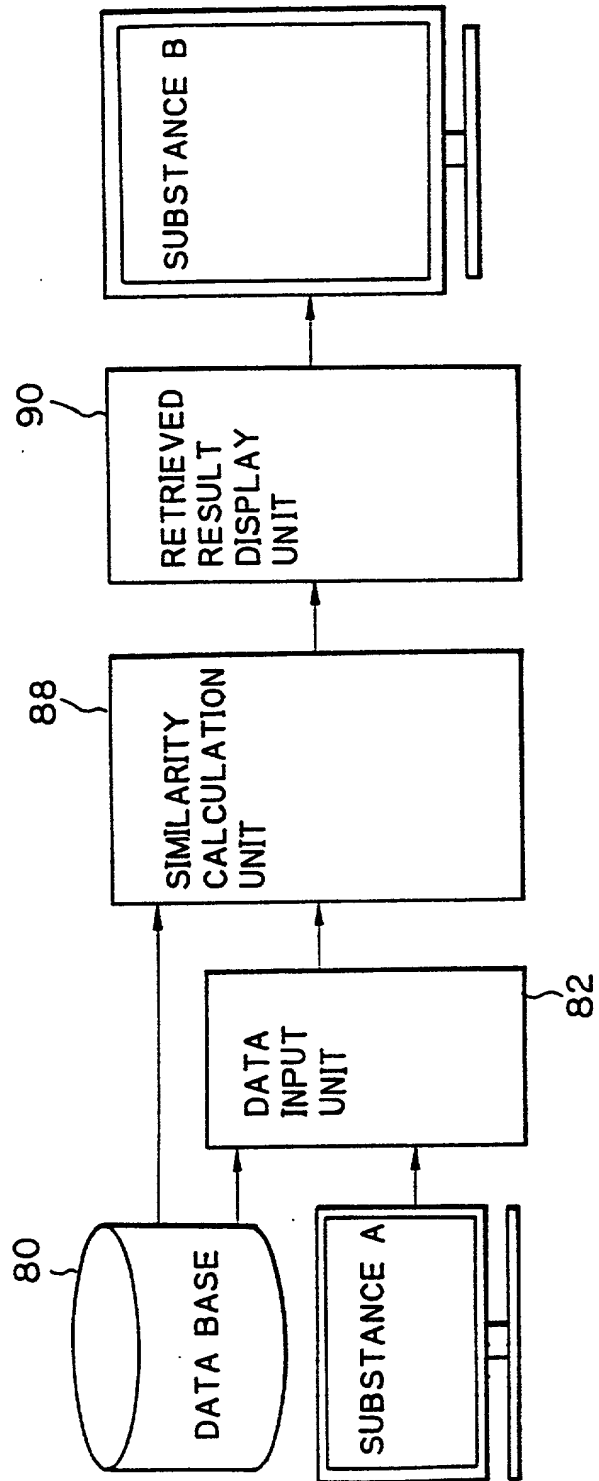


Fig. 28

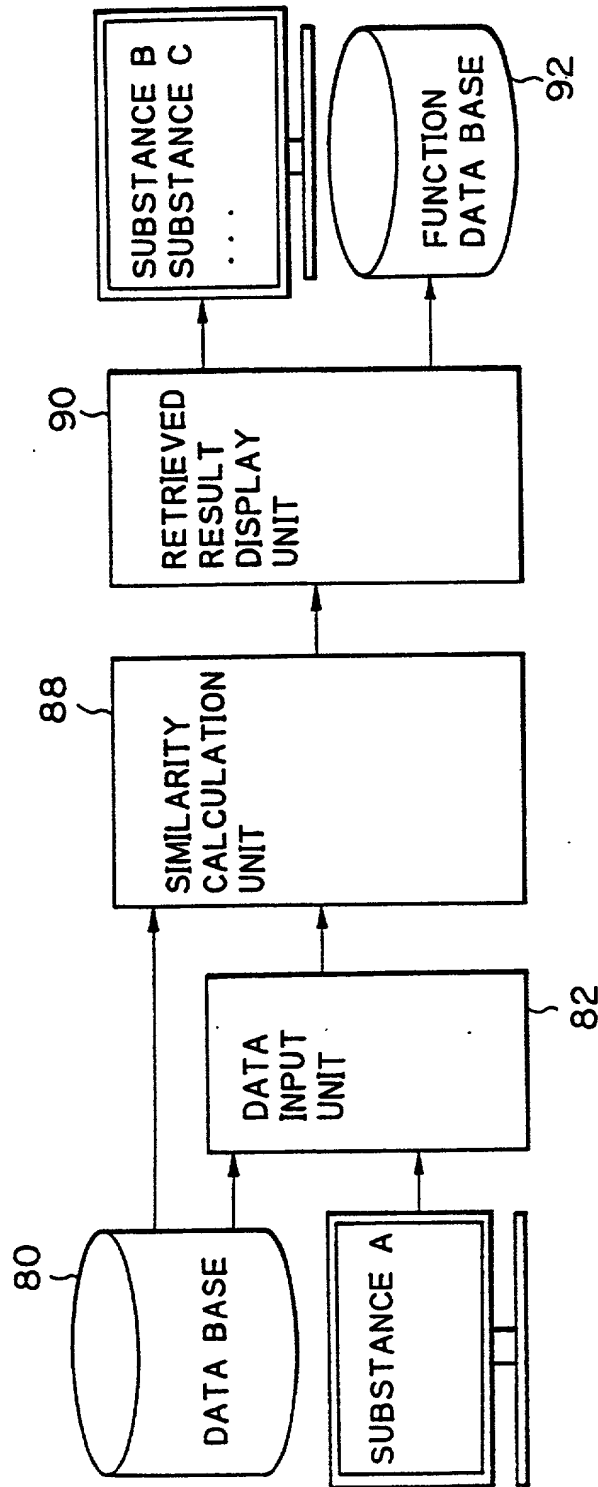


Fig. 29

===== ATP/GTP binding site =====

Probe = (elongation factor)

7 8 9 10 11 12 13 14  
G H V D H G K T < probe >

-----  
8 9 10 11 12 13 14 15  
G A P G S G K G < target >  
G H V D H G K T < probe >  
rmsd=0.648732 adenylylate kinase

unit - A

. : . : .  
10 11 12 13 14 15 16 17  
G A G G V G K S < target >  
G H V D H G K T < probe >  
rmsd=0.421770 ras protein

Fig. 30

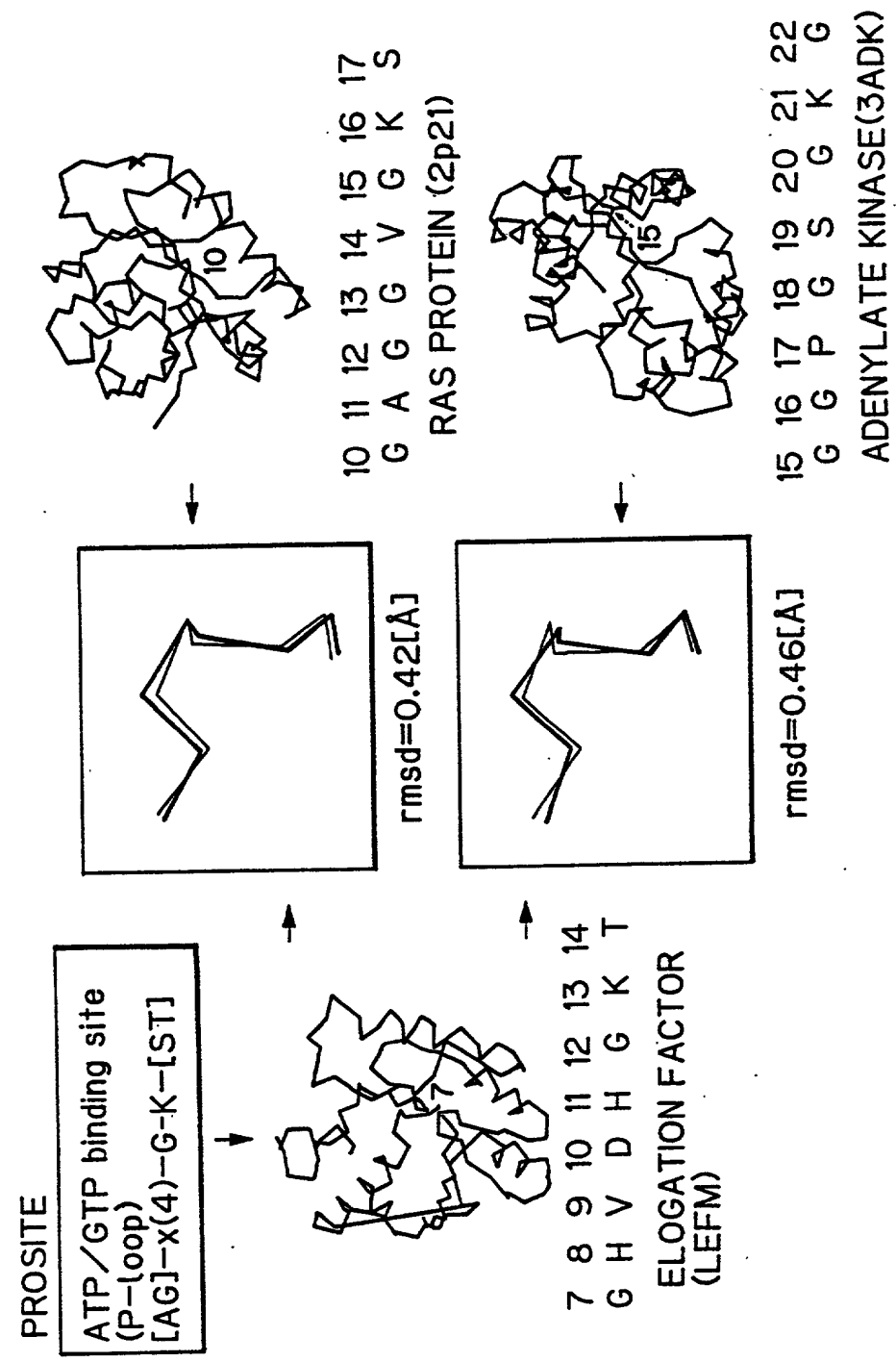


Fig. 31

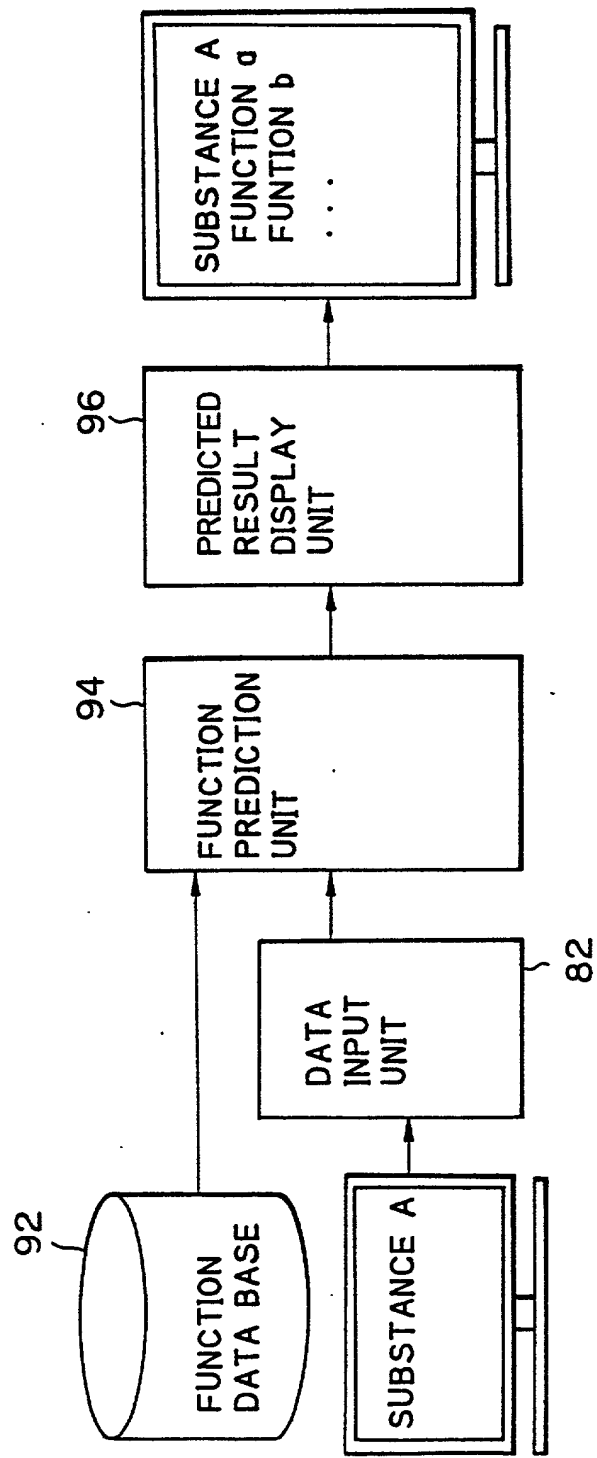


Fig. 32 A

Fig. 32 B

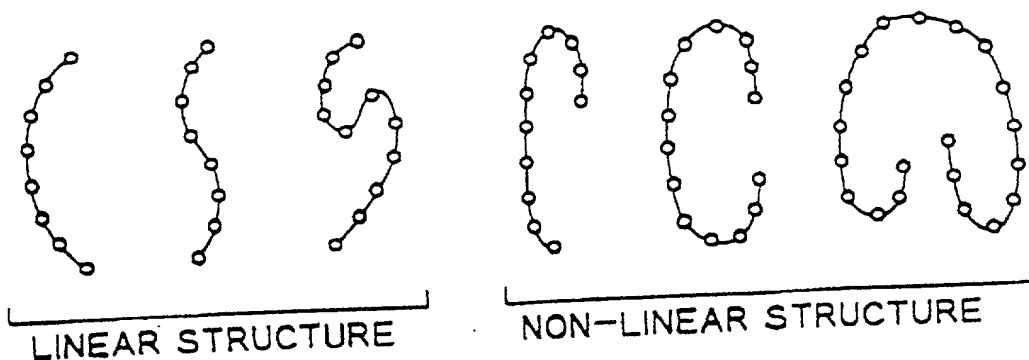
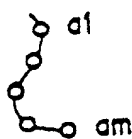
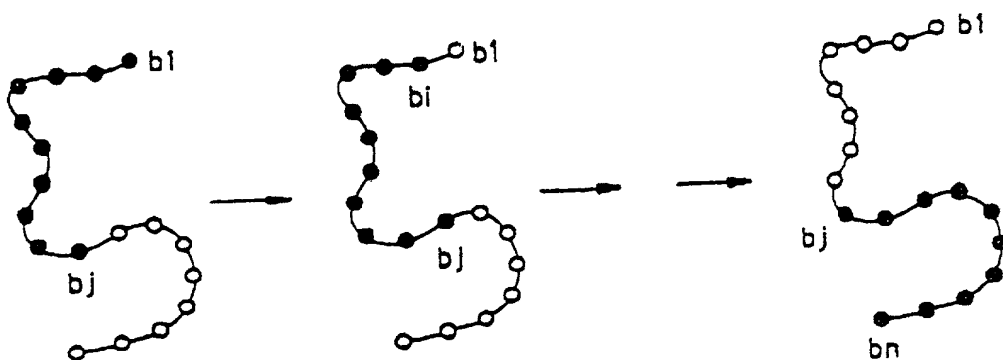


Fig. 33

WHEN  $f(x)=2x$

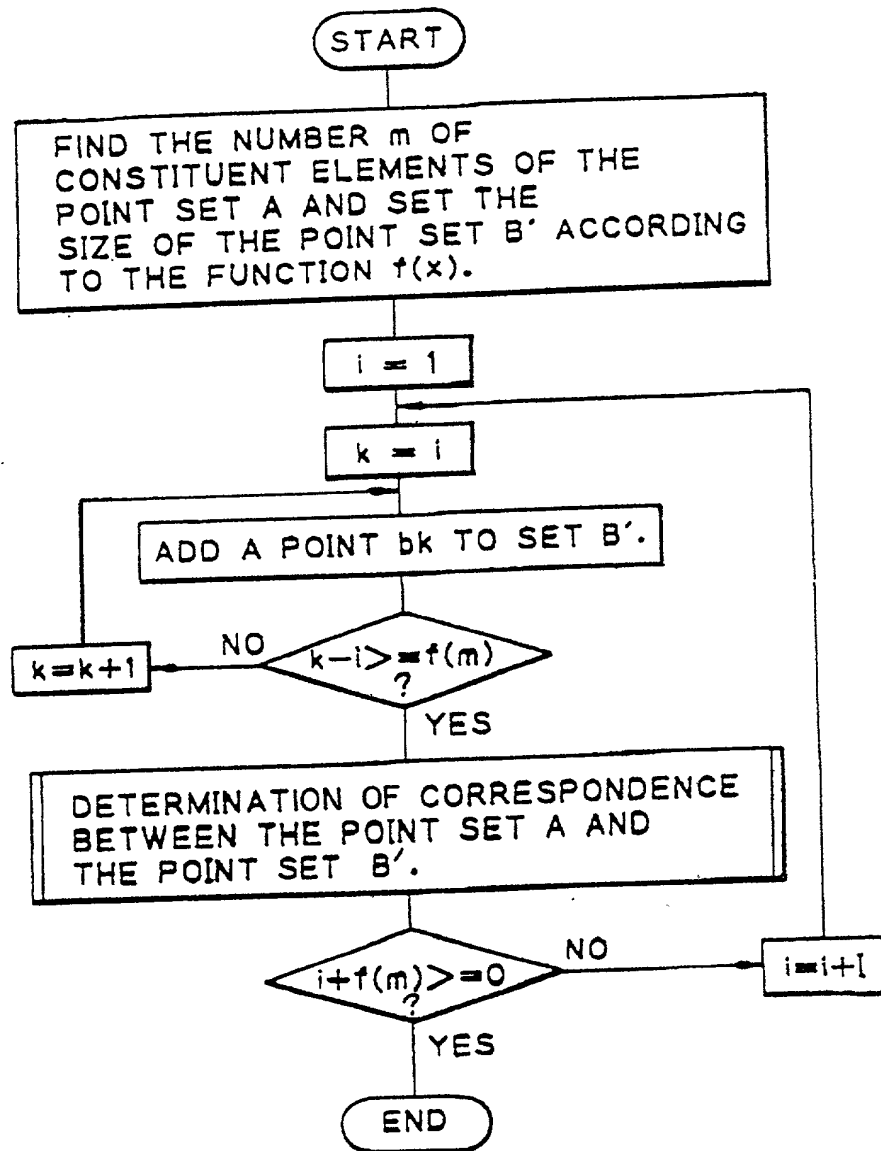


$A=\{a_1, \dots, a_m\}$



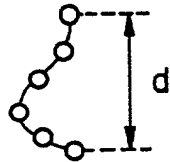
$B=\{b_1, \dots, b_i, \dots, b_j, \dots, b_n\}$

Fig. 34



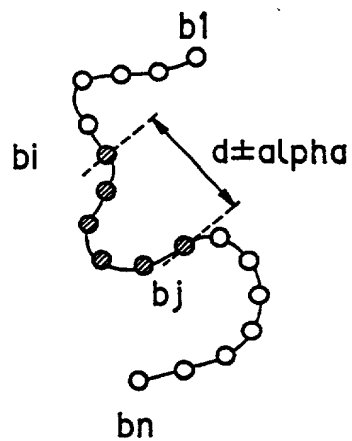


*Fig. 35 A*



$$A = \{a_1, a_2, \dots, a_m\}$$

*Fig. 35 B*



$$B = \{b_1, \dots, b_i, \dots, b_j, \dots, b_n\}$$

Fig. 36

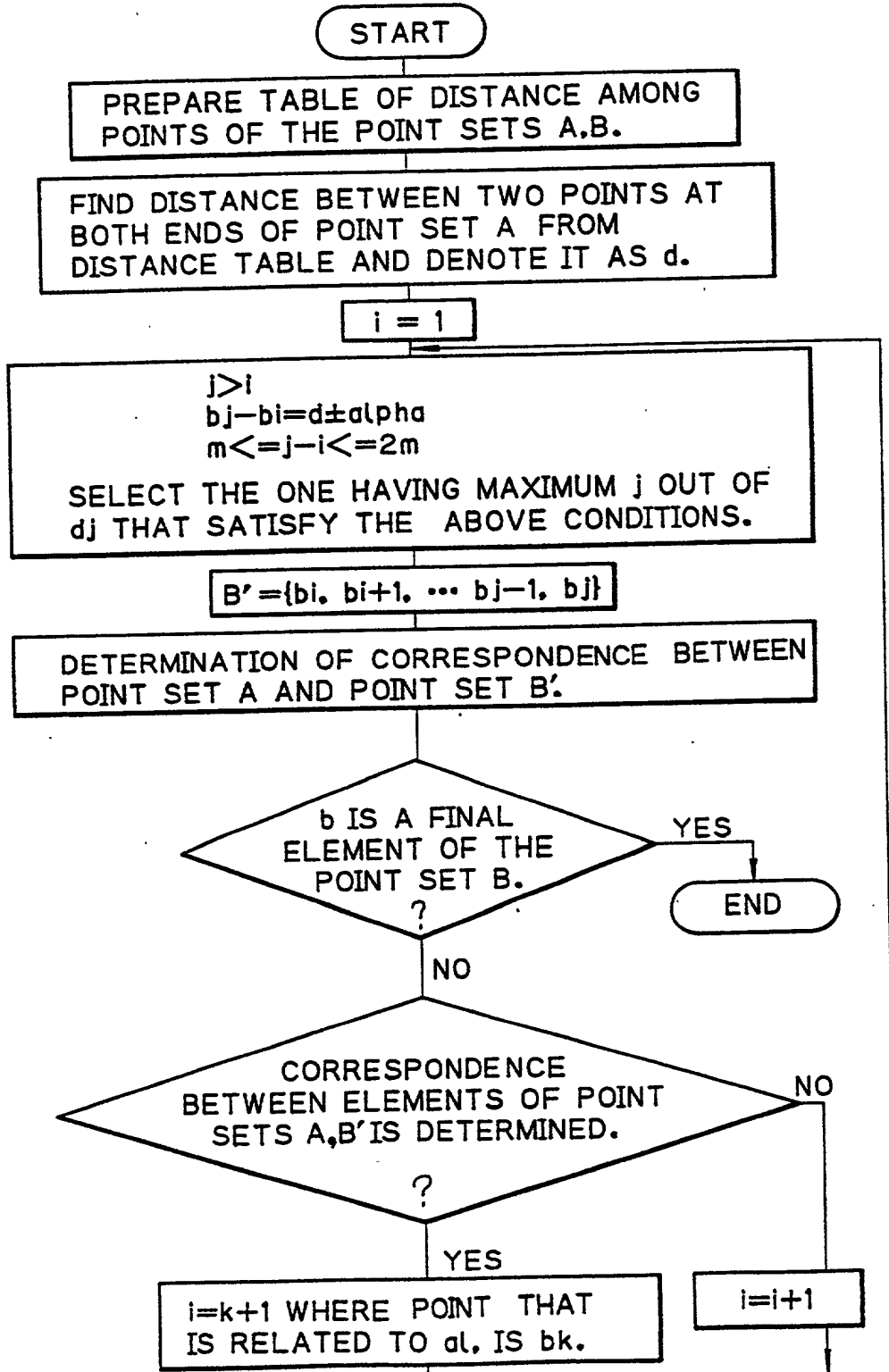
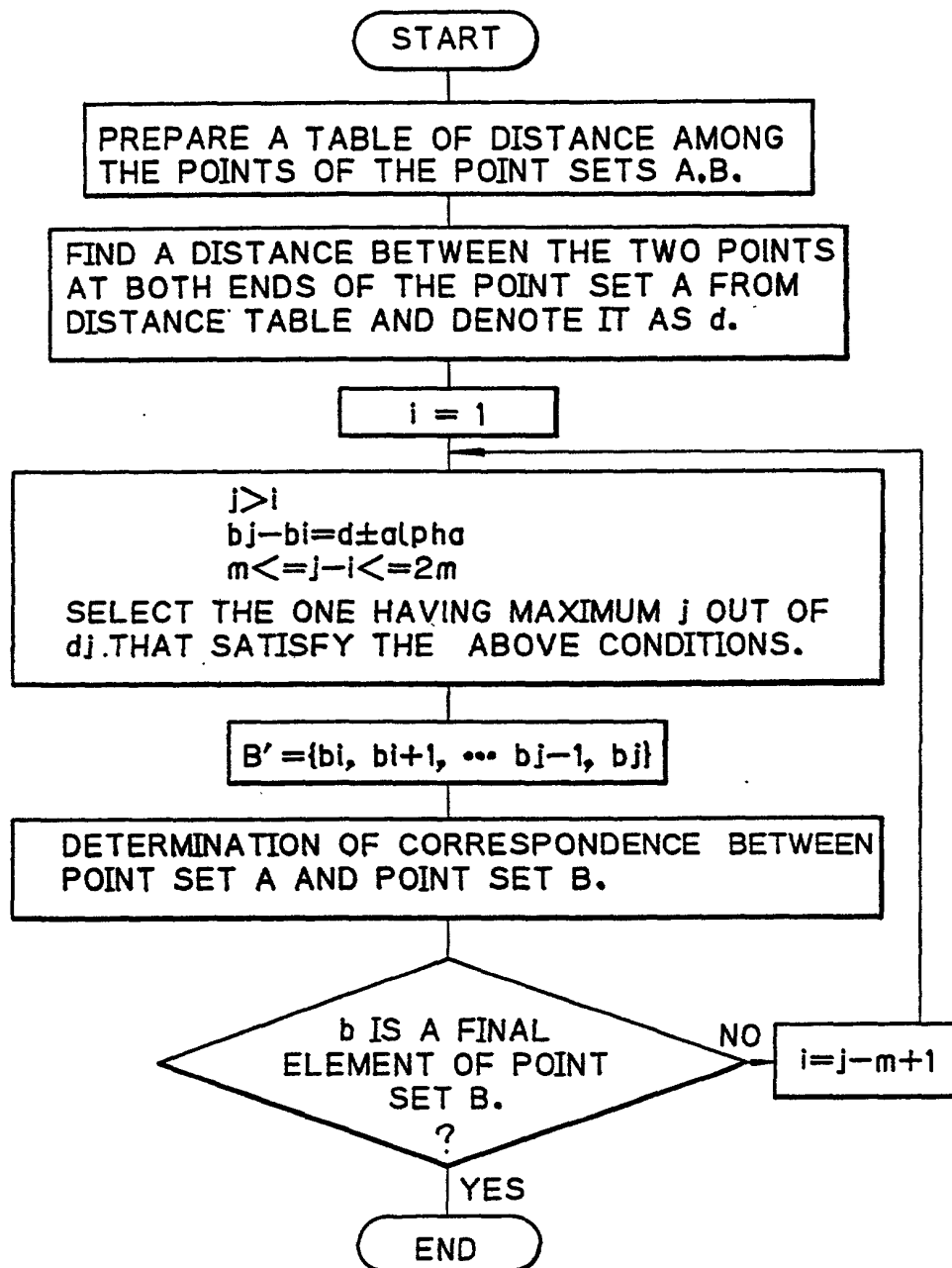


Fig. 37



**Fig. 38 A**

1	I	V	G	G	Y	T	C	C	A	N	T	V	P	Y	Q	V	S	L	N	S
21	G	Y	H	F	C	G	G	S	L	I	N	S	Q	W	V	V	S	A	A	H
41	C	Y	K	S	G	I	Q	V	R	L	G	E	D	N	I	N	V	V	E	G
61	N	E	Q	F	I	S	A	S	K	S	I	V	H	P	S	Y	N	S	N	T
81	L	N	N	D	I	M	L	I	K	L	K	S	A	A	S	L	N	S	R	V
101	A	S	I	S	L	P	T	S	C	A	S	A	G	T	Q	C	L	I	S	G
121	W	G	N	T	K	S	S	G	T	S	Y	P	D	V	L	K	C	L	K	A
141	P	I	L	S	D	S	S	C	K	S	A	Y	P	G	Q	I	T	S	N	M
161	F	C	A	G	Y	L	E	G	G	K	D	S	C	Q	G	D	S	G	G	P
181	V	V	C	S	G	K	L	Q	G	I	V	S	W	G	S	G	C	A	Q	K
201	N	K	P	G	V	Y	T	K	V	C	N	Y	V	S	W	I	K	Q	T	I
221	A	S	N																	

AMINO ACID SEQUENCE OF TRYPSIN (EXCERPT FROM PDB)

**Fig. 38 B**

1	V	V	G	G	T	E	A	Q	R	N	S	W	P	S	Q	I	S	L	Q	Y
21	R	S	G	S	S	W	A	H	T	C	G	G	T	L	I	R	Q	N	W	V
41	M	T	A	A	H	C	V	D	R	E	L	T	F	R	V	V	V	G	E	H
61	N	L	N	Q	N	N	G	T	E	Q	Y	V	G	V	Q	K	I	V	V	
81	P	Y	W	N	T	D	D	V	A	A	G	Y	D	I	A	L	L	R	L	A
101	Q	S	V	T	L	N	S	Y	V	Q	L	G	V	L	P	R	A	G	T	I
121	L	A	N	S	P	C	Y	I	T	T	G	W	G	L	T	R	T	N	G	Q
141	L	A	Q	T	L	Q	Q	A	Y	L	P	T	V	D	Y	A	I	C	S	S
161	S	S	Y	W	G	S	T	V	K	N	S	M	V	C	A	G	G	D	G	V
181	R	S	G	C	Q	G	D	S	G	G	P	L	H	C	L	V	N	G	Q	Y
201	A	V	H	G	V	T	S	F	V	S	R	L	G	C	N	V	T	R	K	P
221	T	V	F	T	R	V	S	A	Y	I	S	W	I	N	N	V	I	A	S	N

AMINO ACID SEQUENCE OF ELASTASE (EXCERPT FROM PDB)

**Fig. 39 A**

Key site number 36 - 41 in Trypsin

41 42 43 44 45 46

M T A A H C < target >

V S A A H C < probe >

d = 12.070038 [A]

r.m.s.d. = 0.061077 [A]

The number of atoms in a probe = 6

The number of atoms in PDB = 240

The number of combination = 1

Time = 1sec

RETRIEVED RESULTS OF HISTIDINE ACTIVE SITES

**Fig. 39 B**

Key site number 175 - 179 in Trypsin

186 187 188 189 190

G D S G G < target >

G D S G G < probe >

d = 8.922721 [A]

r.m.s.d. = 0.092879 [A]

The number of atoms in a probe = 5

The number of atoms in PDB = 240

The number of combination = 1

Time = 1sec

RETRIEVED RESULTS OF SERINE ACTIVE SITES

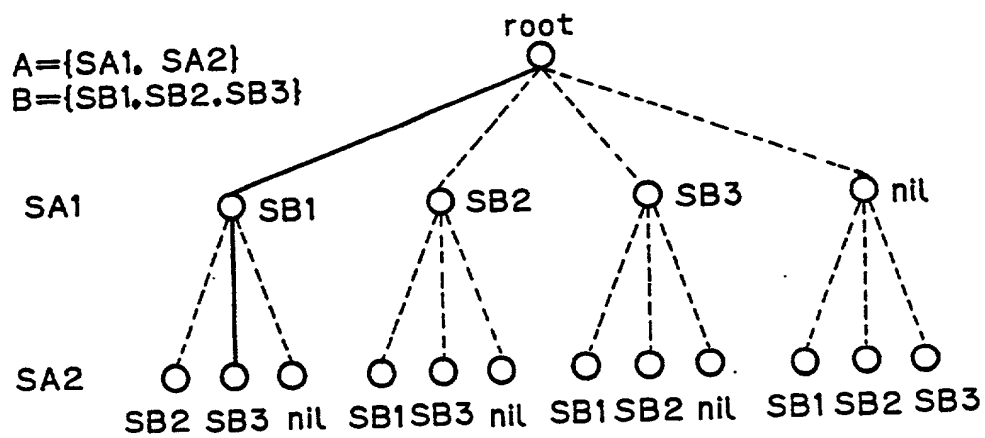
**Fig. 40**

Fig. 41

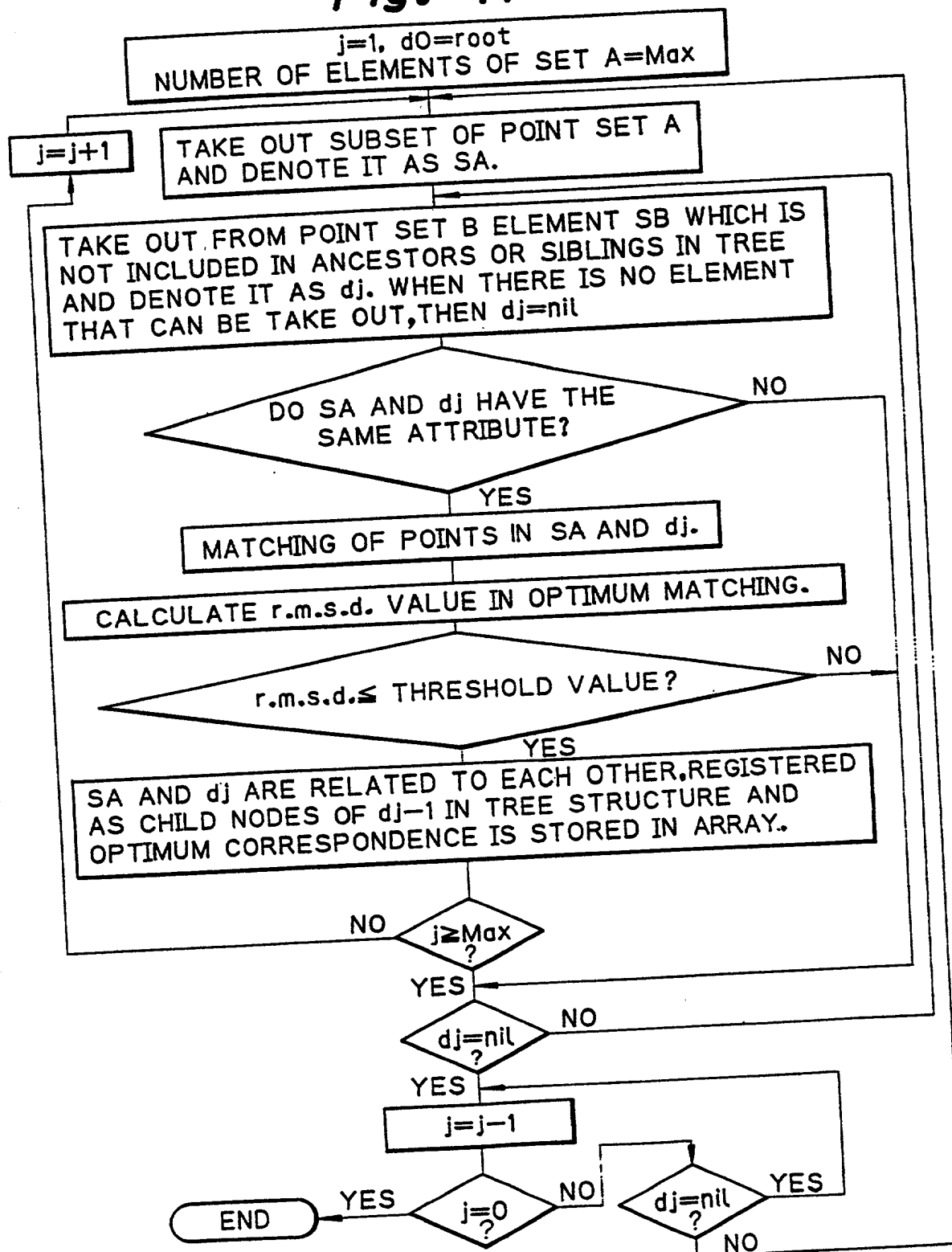


Fig. 42

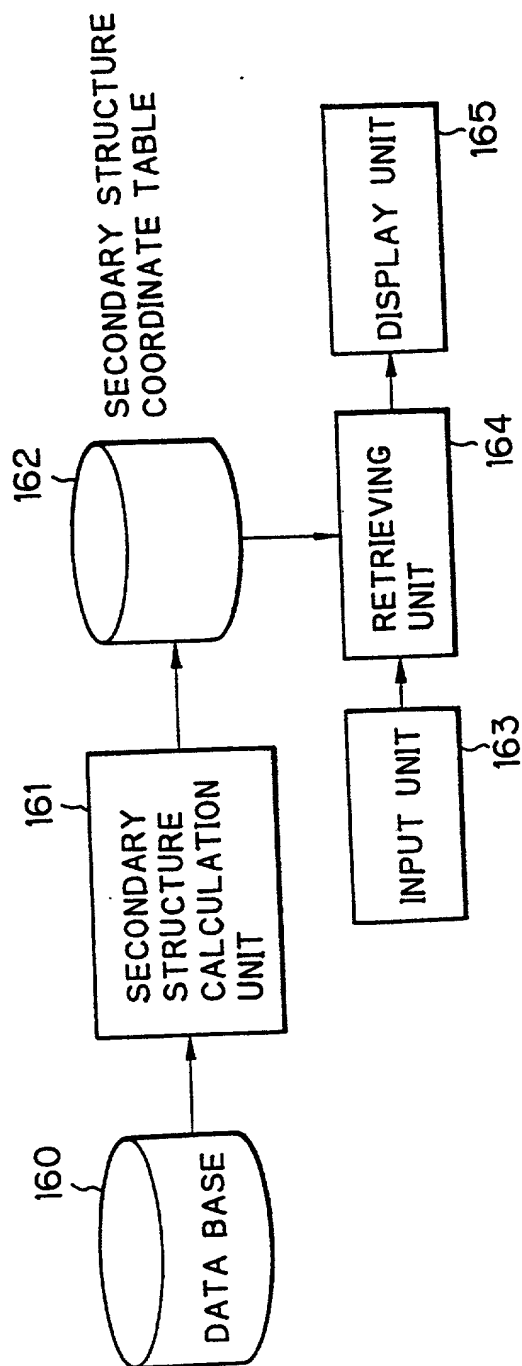
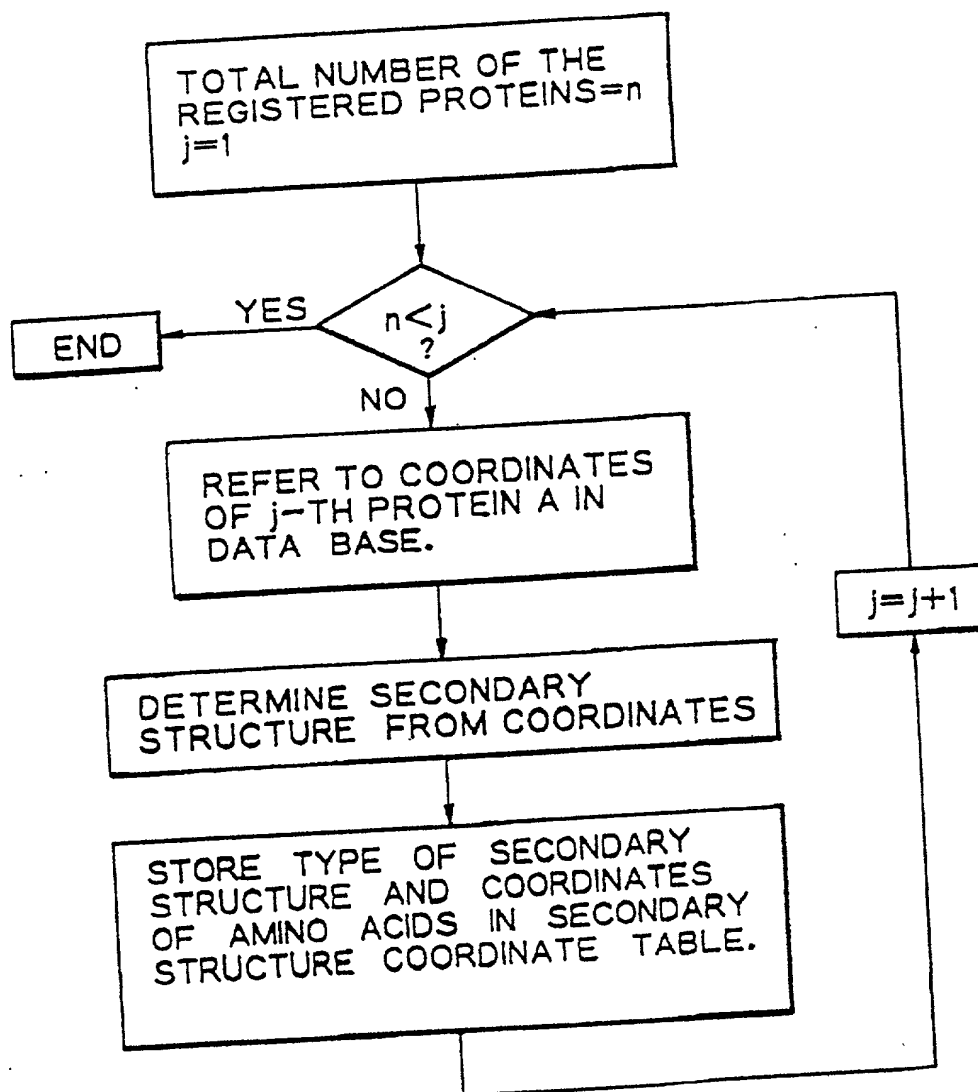




Fig. 43



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SUBSET	COORDINATES	TYPE
S1	{X1,X2,X3,X4,..... Xa}	$\alpha$ - HELIX
S2	{Xa+1,Xa+2,.....Xb}	$\alpha$ - HELIX
S3	{Xb+1,Xb+2,.....Xc}	$\beta$ - SHEET
S4	{Xc+1,Xc+2,.....Xd}	$\beta$ - SHEET
	⋮	⋮
Sn	{Xi+1,Xi+2,.....Xm}	3 - TURN

Fig. 45

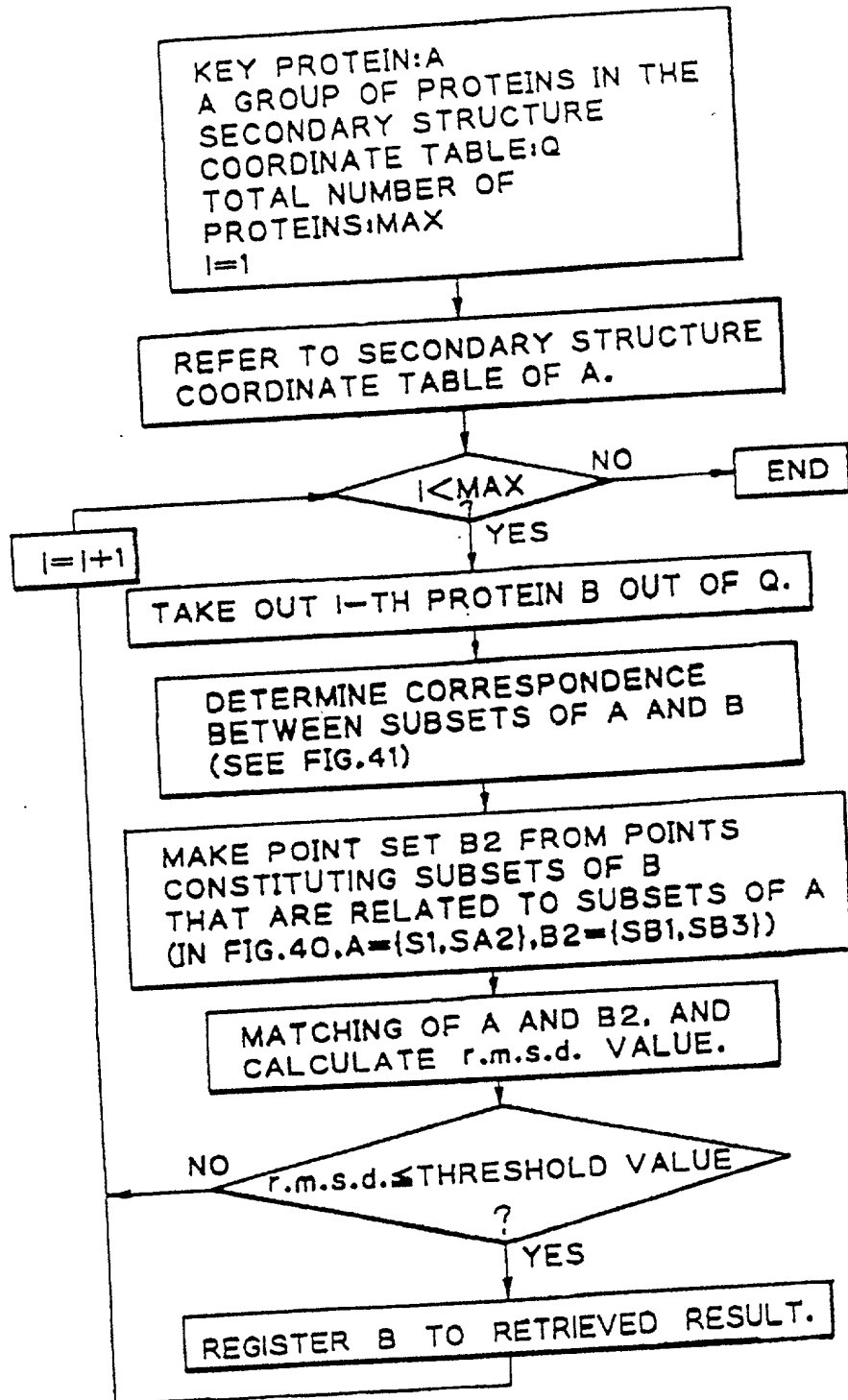


Fig. 46

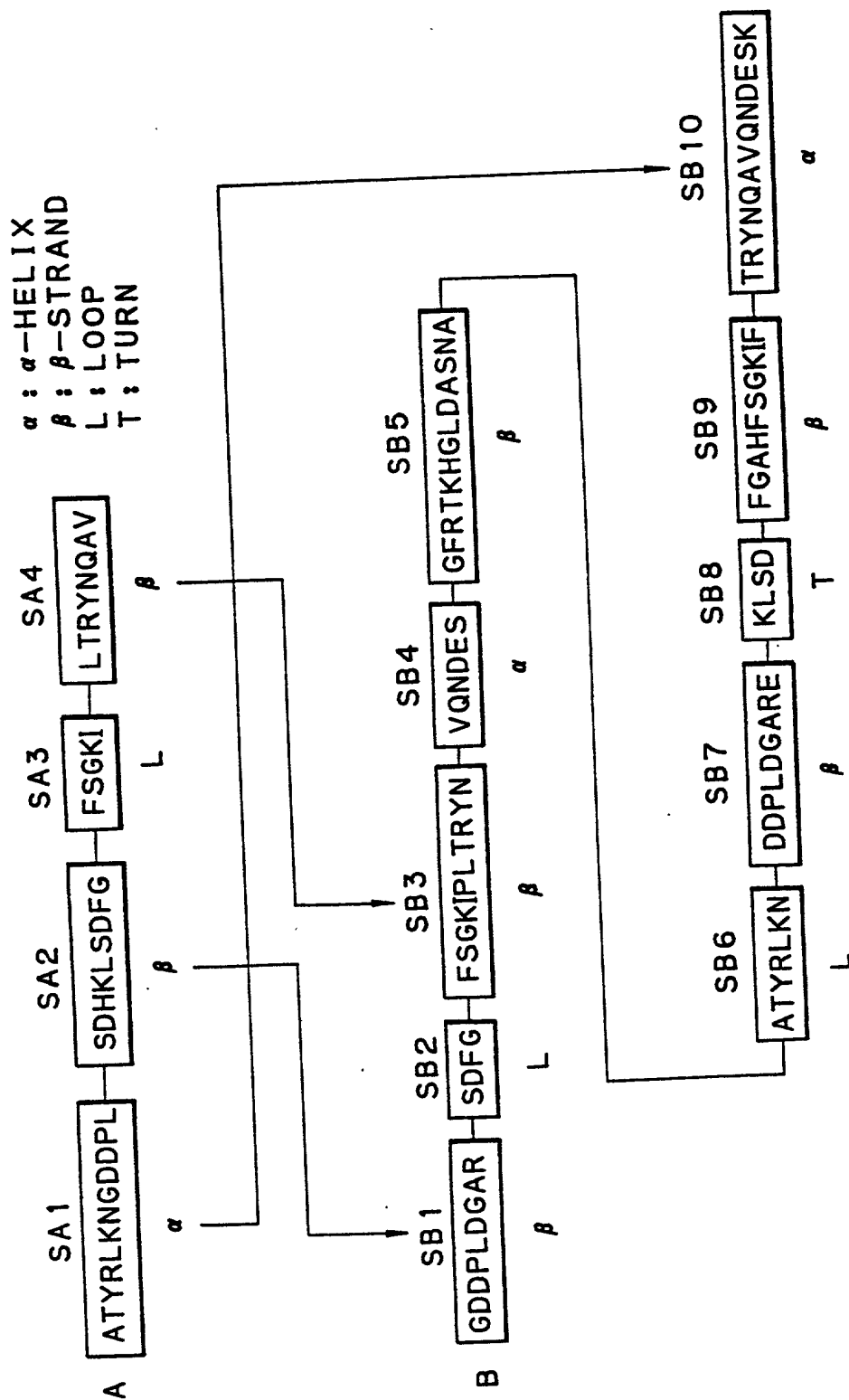
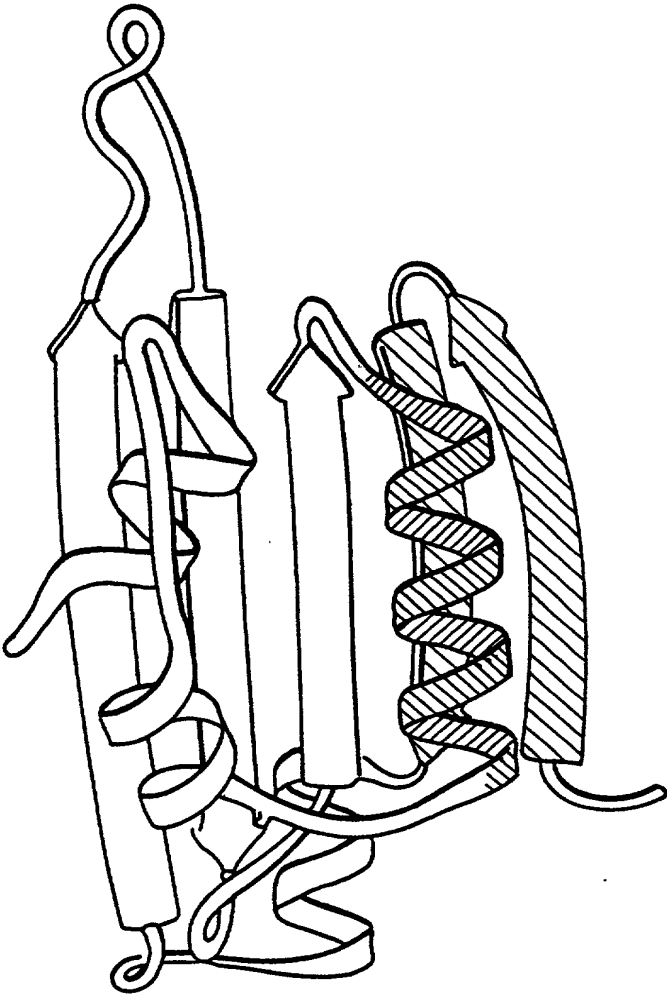
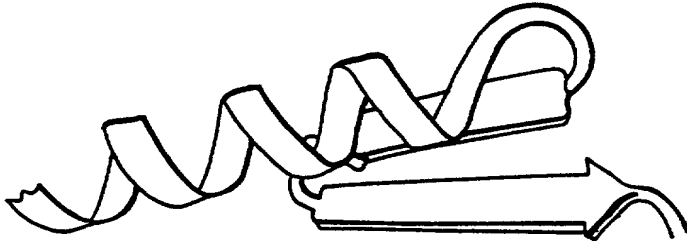


Fig. 47 B



PROTEIN B HAVING A SIMILAR STRUCTURE

Fig. 47 A



KEY PROTEIN A